Minimal code for examples in the main text.

Here we demonstrate the package's usage by presenting a set of minimal R instructions that can reproduce the population dynamics presented in figures 3 and 4. All used functions are either included in MANTIS or present by default in a basic installation of R.

```r
# load package MANTIS
library(MANTIS)

# minimal code for figure 3

# antigenic structure: with 2 loci (6 strains)
epiStruc <- c(3,2)

# get number of strains in the system
nStrains <- extractNumberStrains(epiStruc)

# time at which the solver will stop
tMax <- 415

# solver step
tInt <- 0.005

# record solution every tObsPer steps
tObsPer <- 150

# random initial conditions for infected (Y)
infInitCond <- c(4.307817e-05, 6.552645e-05, 6.619973e-05, 1.698892e-05, 2.037509e-05, 2.093472e-05)

# parameter values here defined by year (output time therefore in years)
beta <- 292
sigma <- (1/5)*365 # 5 days infectious period
mu <- 1/50 # 50 years of life-span

cols <- rev(c("black","purple","blue","limegreen","red","orange"))

# running model with weak cross-immunity
gamma <- 0.1
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

# plot infected class (left, figure 3)
plotY(simdata, x1Obs=0.2, x0Obs=1, yMax=0.0007, colours=cols, yylab="Y", mainRight="NSS")

# running model with intermediate cross-immunity
gamma <- 0.75
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

# plot infected class (centre, figure 3)
plotY(simdata, x1Obs=0.2, yMax=0.015, colours=cols, yylab="Y", mainRight="CSS")

# running model with strong cross-immunity
gamma <- 0.95
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

# plot infected class (right, figure 3)
plotY(simdata, x1Obs=0.2, yMax=0.0007, colours=cols, yylab="Y", mainRight="DSS")

# minimal code for figure 4

define epitope structure, with 2 loci (6 strains)
epiStruc <- c(3,2)
```
#get number of strains in the system
nStrains <- extractNumberStrains(epiStruc)

#time at which the solver will stop
tMax <- 415

#solver step
tInt <- 0.005

#record solution every tObsPer steps
tObsPer <- 150

#random initial conditions for infected
infInitCond <- c(4.307817e-05, 6.552645e-05, 6.619973e-05,
                  1.698892e-05, 2.037509e-05, 2.093472e-05)

#parameter values here defined by year
#thus output time scale will be years
beta <- 292
sigma <- (1/5)*365 # 5 days infectious period
mu <- 1/50 # 50 years of life-span

#running model with weak cross-immunity
gamma <- 0.1
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

#plot infected class (left, black, figure 4)
plotDiversity(simdata, xiObs=0.2, xfObs=1, ymax=2, colour="black", yylab="Y",
mainRight="shannon index, diversity in NSS")

#running model with intermediate cross-immunity
gamma <- 0.75
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

#plot infected class (left, red, figure 4)
plotDiversity(simdata, xiObs=0.2, xfObs=1, ymax=2, colour="red", yylab="Y",
mainRight="shannon index, diversity in CSS")

#running model with strong cross-immunity
gamma <- 0.95
simdata <- runMANTIS(epiStruc, tMax, tObsPer, tInt, infInitCond, beta, gamma, sigma, mu)

#plot infected class (left, blue, figure 4)
plotDiversity(simdata, xiObs=0.2, xfObs=1, ymax=2, colour="blue4", yylab="Y",
mainRight="shannon index, diversity in DSS")

#set a wide range of gammas to calculate single-strain dominance
gammas <- c(0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.75, 0.8, 0.95, 0.99)

#run model for each gamma value and get SSD for each run
y.SSDs <- measureSSDfromGammaRange(gammas, method="host", epiStruc, tMax, tObsPer, tInt, tStep,
infInitCond, beta, sigma, mu, xiObs = 0.7, xfObs = 1)

#plot gammas against SSDs found (centre, figure 4)
plot(gammas, y.SSDs, t='l', col="grey44")

#set a wide range of gammas to calculate strain diversity
gammas <- c(0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.75, 0.8, 0.95, 0.99)

#run model for each gamma value and get Shannon index for each run
y.DIVs <- measureDiversityfromGammaRange(gammas, method="shannon", epiStruc, tMax, tObsPer, tInt, tStep,
infInitCond, beta, sigma, mu, xiObs = 0.7, xfObs = 1)

#plot gammas against DIVs found (right, figure 4)
plot(gammas, y.DIVs, t='l', col="grey44")